**NMR Study of Chain Motion in Atactic** 

Polypropylene at High Pressure

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**ABSTRACT** 

Deuteron solid-state NMR techniques at high pressure are used to study the chain

dynamics in the amorphous polymer atactic polypropylene. The arrest of the structural

relaxation above the glass-transition temperature  $T_{\rm g}$  is investigated using one and two-

dimensional deuteron NMR spectra. The slow reorientation of the main chain segments is

identified with the  $\alpha$ -process observed in mechanical relaxation experiments. On

approaching the glass transition the time scale of the collective motion of the main chain

becomes longer very rapidly at decreasing temperature. Along isobars, at pressure values

up to 5 kbar, the temperature dependence of the logarithmic average correlation time is

very well described by a Vogel-Fulcher-function. The motion of the main chain is

strongly dependent on pressure while its character is mainly determined by the distance to

 $T_{\rm g}$ . The introduction of the equation of state allows the investigation of the dynamic

behavior on isothermal and isochoric paths on approaching  $T_g$ . It is found that along an

isothermal the mobility as a function of the density is also of the Vogel-Fulcher form.

**KEYWORDS:** atactic polypropylene, glass-transition, high pressure, NMR, slow chain

motion

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## 1. INTRODUCTION

The arrest of the structural relaxation at the glass transition is an important aspect of polymer glasses. At temperature values just above the glass transition the diffusive motion and the conformational changes of the main chain (the  $\alpha$ -process) slow down drastically. In amorphous polymers its temperature dependence is very well described by the Williams-Landel-Ferry equation<sup>1,2</sup> or, equivalently, by the Vogel-Fulcher-Tammann-Hesse function<sup>3,4,5</sup>. The time scale of the structural relaxation changes over about fifteen orders of magnitude. The large change in the mechanical properties at the glass transition is related to the change in the molecular dynamics.

In this paper we present a study of the effect of high pressure on the chain dynamics of atactic polypropylene (aPP) in the liquid close to the glass transition. This study is an extension to conditions of high pressure of the work in deuterated aPP by Spiess and coworkers<sup>6,7</sup> at ambient pressure with the technique of two-dimensional exchange deuteron NMR. They found that the spectra obtained could be successfully reproduced by assuming isotropic rotational diffusion as the model for the slow reorientation of the main chain segments. All data obtained from these experiments, combined with those determined earlier through measurements of <sup>13</sup>C spin-lattice relaxation times<sup>8</sup> at elevated temperatures could be represented quantitatively over more than twelve orders of magnitude by the WLF equation. The conclusion was that the time scales of the chain dynamics as probed for individual chain segments by 2D NMR are in excellent agreement with those of the collective processes responsible for the strong temperature dependence of the  $\alpha$ -relaxation process.

In this study we first will show that at high pressure, up to 5 kbar, the slowing down of the main chain motion on approaching the glass transition can be represented very well by the Williams-Landel-Ferry equation. Next, we will introduce the equation of state to investigate the dynamic behavior on isothermal and isochoric paths, in order to separate the effects of the changes in temperature and in density on approaching the glass transition.

## 2. EXPERIMENTAL

We used a sample of 88% uniformly deuterated atactic polypropylene, prepared from the same material as was used in ref<sup>6</sup>. The molecular weight (by viscometry) is  $M_w \approx 25000$  g mol<sup>-1</sup>. With differential scanning calorimetry the value of  $T_g$  of this

deuterated aPP material was determined to be 253 K at ambient pressure.

The NMR spectrometer and high-pressure NMR probe used for the investigation presented in this paper has been described in detail elsewhere<sup>9</sup>. The experiments have been performed at a  $^2$ H-NMR frequency of 41.433 MHz with  $\pi/2$  radio frequency pulses of 4  $\mu$ s.

We obtained one-dimensional quadrupole echo<sup>10</sup> spectra using the standard sequence of two radio-frequency pulses with a phase difference  $\pi/2$ , separated by a time interval  $\tau_I$ . We obtained two-dimensional exchange spectra by using the five-pulse sequence described in reference<sup>11</sup>.

High-purity (at least 99.999 %) helium gas is used as the pressurizing medium. Helium is used because of its very small solubility in hydrocarbon liquids and elastomers. The equipment for the generation and measurement of pressure, has also been described in reference<sup>9</sup>.

## 3. NMR SPECTRA

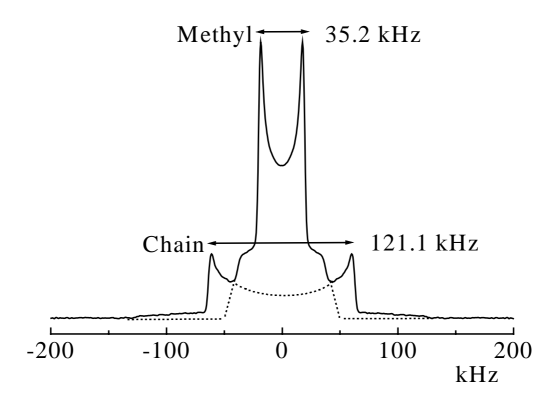
## 3.1. <sup>2</sup>H-NMR quadrupole echo spectra

The interaction dominating the  ${}^{2}$ H-NMR spectra in deuterated hydrocarbon polymers is the coupling between the electric quadrupole moment eQ of the deuteron (with spin quantum number I=1) and the electric field gradient (EFG) at the site of the deuteron. For a single C-D bond, the combined Zeeman and quadrupole Hamiltonians lead to a deuteron NMR spectrum consisting of only two lines, at frequencies:

$$\omega = \omega_0 \pm \delta_Q \left( 3\cos^2 \theta - 1 + \eta \sin^2 \theta \cos 2\varphi \right) = \omega_0 \pm \omega_Q (\theta, \varphi)$$
 (1)

where  $\omega_0$  is the deuteron Zeeman frequency and  $\delta_Q \equiv 3e^2qQ/8\hbar$ , while eq is the value of the principle value  $\partial^2 V/\partial z^2$  of the EFG tensor;  $\eta$  is the asymmetry parameter  $(\partial^2 V/\partial x^2 - \partial^2 V/\partial y^2)/\partial^2 V/\partial z^2$ . The quadrupole frequency shift  $\omega_Q$  depends on the polar angles  $\theta$  and  $\varphi$  of the static magnetic field  $\mathbf{B}$  in the principle axes system (PAS) of the EFG tensor. The much weaker magnetic dipole-dipole interactions, present in the system of nuclear spins, result in a homogeneous broadening (to a width of only about 1 kHz) of each line. In an aliphatic C-D bond the EFG is nearly axially symmetric ( $\eta$ =0) and the quadrupole constant  $e^2qQ/\hbar$  is  $2\pi\cdot 169\cdot 10^3$  rads<sup>-1</sup>.

For an isotropic distribution of C-D bond orientations, such as present in a rigid amorphous deuterated-polymer solid, the quadrupole echo spectrum arising from the two



**Fig. 1.** The experimental quadrupole echo <sup>2</sup>H NMR spectrum of aPP in the glassy state at 246 K and 1 bar. The inner and outer Pake-doublets originating from methyl and chain deuterons, respectively, are indicated schematically.

transitions in equation (1) is the Pake doublet<sup>12</sup>, with characteristic singularities at frequencies  $\pm \delta_Q$  corresponding to  $\theta = \pi/2$ . Fig. 1 shows the one-dimensional deuteron quadrupole echo spectrum in the glassy state at ambient pressure at 246 K. As is shown schematically, the spectrum consists of a superposition of two Pake doublets. The broad component with a distance between the singularities  $\Delta_{\text{chain}}$  of about 123 kHz originates from deuterons directly bonded to the almost immobile main chain. The distance is slightly smaller than the value 127 kHz expected from the value of the quadrupole coupling constant. This is due to restricted reorientation of the main chain C-D bonds, occurring on a time scale of  $10^{-6}$  s, shorter than  $\delta_Q^{-1}$ . The narrow component with a distance between the singularities  $\Delta_{\text{methyl}}$  of about 37 kHz is the contribution from the deuterons of the methyl groups, which perform rapid anisotropic reorientation on a time scale of about  $10^{-10}$  s, very much shorter than  $\delta_Q^{-1}$ . As a result, the methyl deuterons experience an average EFG which is axial about the  $C_3$  axis with  $\overline{eq} = eq/3$  for a tetrahedral configuration of bonds.

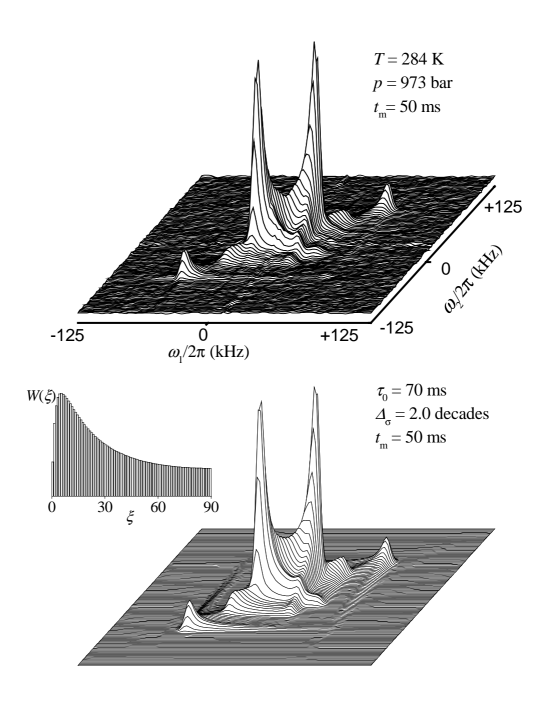
## 3.2. <sup>2</sup>H 2D exchange NMR spectra

For the observation of the very slow motion occurring just above the glass transition temperature, on a time scale of  $10^{-5}$  to  $10^2$  s, we made use of the technique of deuteron two-dimensional (2D) exchange NMR. Theory and application of this technique have been comprehensively presented in the textbook by Schmidt-Rohr and Spiess<sup>13</sup>. As mentioned above, we obtained 2D exchange spectra by using the five-pulse sequence described in reference<sup>11</sup>.

The 2D exchange absorption spectrum  $S(\omega_1, \omega_2; t_m)$  may be identified with the joint probability density  $P(\omega_1, \omega_2; t_m)$  that a nucleus has an NMR frequency  $\omega_1$  before and a frequency  $\omega_2$  after the "mixing" time  $t_m$  included in the pulse sequence. As has been discussed above, the <sup>2</sup>H-NMR frequency is a function of the orientation of the external magnetic field  $B_0$  with respect to the principle axis system of the EFG present at the site of the nucleus. Since in a  $C^2H$  bond the EFG is approximately axially symmetric about the bond direction, one can map the frequency domain  $(\omega_1, \omega_2)$  onto the polar angle domain  $(\theta_1, \theta_2)$  and associate with  $P(\omega_1, \omega_2; t_m)$  the joint probability density  $P'(\Theta_1, \Theta_2; t_m)$  of finding the angles  $\Theta_1$  and  $\Theta_2$  between the  $C^2H$  bond and  $B_0$  before and after a time interval  $t_m$ , respectively. From this one can construct the distribution  $W(\xi, t_m)$  of the angle  $\xi$  of reorientation of the  $C^2H$  bond after a time interval  $t_m$ . The angular information contained in a <sup>2</sup>H 2D exchange spectrum is completely specified by the one-dimensional reorientation angle distribution  $W(\xi, t_m)$ . By systematically varying  $t_m$ , one is able to monitor the dynamic evolution of  $P(\omega_1, \omega_2; t_m)$  and of  $W(\xi, t_m)$ , and to determine the type and time scale of the molecular reorientation.

The above is only strictly true in the so-called slow motion-limit, in which a  $C^{-2}H$  bond reorients only during  $t_m$  and not in  $t_1$ , nor in  $t_2$ . The interpretation given is therefore only valid in practice for motion with a characteristic time longer than  $10^{-3}$  s. In the intermediate exchange regime  $(10^{-5} \text{ to } 10^{-3} \text{ s})$ , the correspondence between the 2D exchange spectrum  $S(\omega_1, \omega_2; t_m)$  and the joint probability density  $P(\omega_1, \omega_2; t_m)$  breaks down. However, by incorporating the effect of motion on an intermediate time scale on the calculated 2D spectra<sup>7,14</sup>, the dynamic range for which experimental and calculated spectra can be compared and reorientation angle distributions  $W(\xi; t_m)$  are obtained, is extended from  $10^{-3}$  to  $10^{-5}$  s.

Fig. 2 shows a typical <sup>2</sup>H 2D exchange spectrum  $S(\omega_1, \omega_2; t_m)$  of aPP. It was determined



**Fig. 2.** Typical <sup>2</sup>H 2D exchange spectrum of aPP. (a) Experiment at 973 bar and 284 K with a mixing time  $t_m$ =50 ms. (b) Calculated spectrum with  $\tau_0$ =70 ms and  $\Delta_{\sigma}$ =2 decades. The inset shows the corresponding reorientation angle distribution W( $\xi$ ).

at 973 bar and at 284 K, which is 17 K above  $T_g$  at that pressure. The mixing time  $t_m$ =50 ms. The spectrum appears to be very similar to the one shown in reference<sup>6</sup>, determined at ambient pressure at 270 K, which is also 17 K above  $T_g$ .

Along the  $\omega_1$ = $\omega_2$  diagonal we recognize the shape of the 1D  $^2$ H spectrum in Fig. 1, namely a superposition of the two Pake patterns originating from the chain and the methyl deuterons. The intensity of the spectrum along the diagonal represents  $C^{-2}H$  bonds which after 50 ms are found to have the same angle with respect to the magnetic field  $B_0$  as at the start of the experiment. Off-diagonal intensity, in particular the ridges parallel to the singularities in  $\omega_1$  and in  $\omega_2$ , indicates reorientation motion occurring at the time scale of the mixing time. The reorientation of the main chain segments causes exchange within both the chain and the methyl deuteron frequencies. The exchange is visible covering all frequencies within two square-like areas. This can be understood as a superposition of 2D-spectra according to a distribution of reorientation angles. The ridges  $S(\pm \delta_Q, \omega_2)$  at  $\omega_1 = \pm \delta_Q = \pm 18$  kHz (the peaks of the methyl Pake pattern) correspond to C-C<sup>2</sup>H<sub>3</sub> bonds that have an orientation perpendicular to B at the start of the mixing period. From these ridges it can already be seen that the reorientation angle distribution  $W(\xi;t_{\rm m})$  is a continuous distribution, indicating that after 50 ms the axis of the methyl group has a finite probability to be found in any orientation with respect to the magnetic field, regardless of its starting position.

Reorientation over discrete and fixed angles would cause typical elliptical ridge patterns<sup>13</sup>. These are not visible in our 2D spectra of aPP.

We performed  $^2$ H 2D exchange NMR experiments along isobars at 1, 973, 1978 and 4986 bar, in temperature ranges from about 11 to about 34 K above  $T_{\rm g}$ . At each temperature and pressure value we obtained two 2D absorption spectra by choosing two different mixing times  $t_{\rm m}$ , with a ratio of 3 to 10. At the lower end of each temperature range the correlation time becomes much longer than the mixing time, allowed by the spin-lattice relaxation time.

We used the values of  $T_g$  at each pressure value as they have been obtained from the decay of the quadrupole echo in reference<sup>9</sup>, where the data could be represented by the quadratic function:

$$p(T_g) = c_1 (T_g - T_g^0) + c_2 (T_g - T_g^0)^2$$
 (2)

in which  $T_g^{\ 0}$  is the glass-transition temperature at zero pressure, with the following values of the parameters:  $c_1$ =55(±12) barK<sup>-1</sup>,  $c_2$ =0.48(±0.16) barK<sup>-2</sup> and  $T_g^{\ 0}$ =251.2(±2.1) K.

#### 4. THE MODEL FOR THE CHAIN REORIENTATION

In the analysis of the 2D-exchange spectra we assume that in aPP the reorientation motion of the chain segments above the glass transition temperature results in isotropic rotational diffusion of the directions of both the main chain C-<sup>2</sup>H bonds and the C-C<sup>2</sup>H<sub>3</sub> bonds, with the same diffusion coefficient  $D_r$ . This process can also be characterized by a correlation time  $\tau_d$ =1/6 $D_r$ . Spiess and coworkers have shown (reference<sup>15</sup> equations (6) and (10)), that, essentially, the 2D absorption spectrum  $S(\omega_1, \omega_2; t_m)$  can be derived from the rotational diffusion equation.

The characterization of the time scale of the motion requires the introduction of a distribution of correlation times. This is immediately clear, for instance, in the experimental spectrum at 1 bar and 284 K of Fig. 3, where a narrow peak indicating fast quasi-isotropic motion occurs in the center, together with the effects of exchange on a much slower time scale.

The occurrence of a correlation time distribution is related to the non-exponential loss of correlation in the chain relaxation process<sup>13</sup>, following the Kohlrausch-Williams-Watts (KWW) relaxation function<sup>16</sup>  $C(t)=\exp[-(t/\tau_{\rm KWW})^{\beta}]$ . The correlation time distribution, obtained as the Laplace transform of the KWW function, resembles a Gaussian distribution of  $\ln \tau$  values, apart from showing a more rapid decrease in the wing at long  $\tau$  values.

Although possible, in principle, to make the data analysis by using correlation time distributions, obtained as Laplace transforms of KWW functions with varying  $\tau_{KWW}$  and  $\beta$  values, we choose the simpler approach of using a Gaussian distribution of  $\ln \tau_{d}$  values:

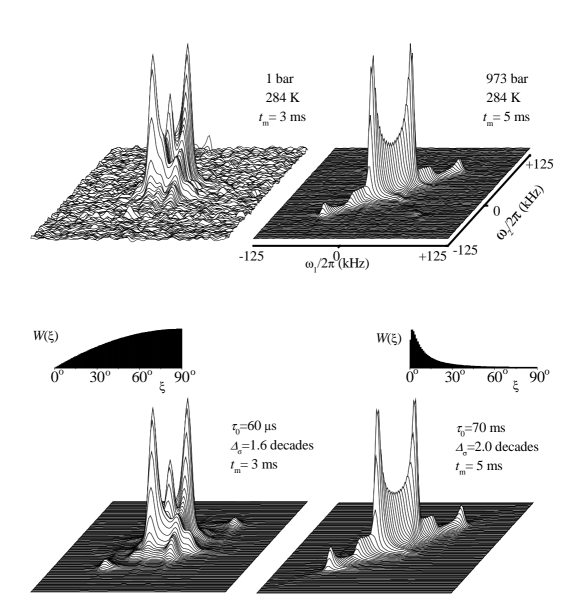
$$\rho(\ln \tau_d) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left\{-\frac{(\ln \tau_d - \ln \tau_0)^2}{2\sigma^2}\right\}$$
(3)

where  $\ln \tau_0$  is the mean value of  $\ln \tau_d$  and  $\sigma$  is the standard deviation of the distribution. The width  $\Delta_{\sigma}$  of the distribution expressed in decades is  $2\sigma \cdot {}^{10}\log(e)$ .

## 4.1. The calculation of spectra

We compared the experimental spectra with spectra calculated using a computer program based on the analysis as presented in references<sup>7,15,14</sup>. 2D time domain responses generated by a five-pulse sequence are calculated, using rotational diffusion

as the model



**Fig. 3.**  $^{2}$ H 2D exchange spectra as a function of pressure. The experiments were performed at 1 bar and 284 K and at 973 bar and 284 K with mixing times  $t_{m}$ =3 ms and 5 ms, respectively.

for the chain segment reorientation. This calculation is performed separately for the chain deuterons and for the methyl deuterons, using axially symmetric EFG tensors. We

account for the effect of the fast small angle reorientation, resulting in the reduced distance of the Pake singularities in the 1D spectra by using reduced quadrupole coupling constants  $2\pi \cdot 161.6 \cdot 10^3$  and  $2\pi \cdot 48.3 \cdot 10^3$  rad s<sup>-1</sup> for the chain deuterons and the methyl deuterons, respectively. In this way the Pake singularities in the calculated spectra appear at the frequencies of the maxima in the experimental spectra.

As is shown in Fig. 2, the experimental spectrum at 973 bar and 284 K obtained with  $t_{\rm m}$ =50 ms, can be reproduced very well by choosing a correlation time distribution with  $\tau_0$  = 70 ms and  $\Delta_{\rm G}$  = 2 decades. Also shown is the reorientation angle distribution  $W(\xi)$  corresponding to this distribution of  $\tau_{\rm d}$ . Another experiment with  $t_{\rm m}$ =5 ms was performed at the same temperature and pressure (see Fig. 3), and an excellent fit is obtained using a distribution with precisely the same parameters.

A clear example of the effect of pressure on the dynamics is shown in Fig. 3. Here we compare two spectra at 284 K, at 1 bar and at 973 bar. The experiments have been performed with approximately equal mixing times, namely  $t_m$ =3 ms and 5 ms, respectively. At 1 bar a lot of exchange intensity is visible, as is the peak in the center of the spectrum due to close to isotropic motion, indicating that a substantial part of the main chain segments are reorienting with correlation times  $\tau_d \le 1$   $\mu$ s. After an increase of the pressure to about 1 kbar the exchange intensity almost disappears. This can also be seen from the reorientation angle distributions and the correlation time distributions, as obtained from fitting calculated spectra to the experimental spectra. The logarithmic average correlation time  $\tau_0$  decreases three orders of magnitude on increasing the pressure to about 1 kbar.

The use of 2D exchange NMR to observe molecular reorientation is only possible within a limited temperature range. At the lowest temperature the spin-lattice relaxation time limits the mixing time and, in the case of aPP, a correlation time  $\tau_d$  longer than approximately 100 s cannot be observed. At high temperatures the spectrum collapses into an isotropic peak for  $\tau_d \approx 1$   $\mu$ s. In practice, this means that for  $\tau_0 \leq 50$   $\mu$ s, for aPP corresponding to temperatures  $T \geq T_g + 35$  K, information about the molecular reorientation cannot be obtained any more from 2D exchange spectra.

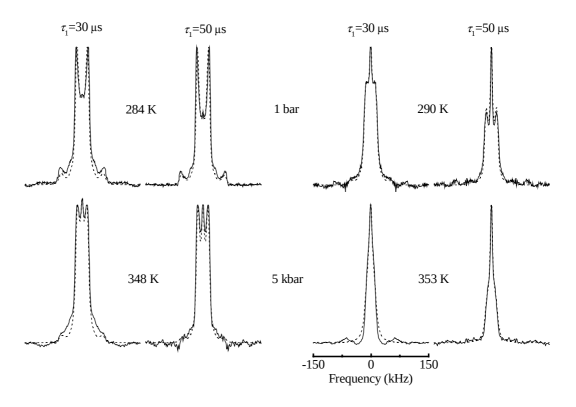
Of course, the motion of the chain segments cannot truly be isotropic, because of the restrictions imposed on the chain segments by the polymer main chain itself and by its environment. We note that the quadrupole coupling Hamiltonian possesses inversion symmetry, reorientation angles  $\xi$  cannot be distinguished from angles  $\pi-\xi$ . Therefore,

quasi-isotropic motion of the chain segments will still result in the same line shapes as produced by isotropic motion.

It has to be noted that, as a 2D exchange spectrum only correlates the orientations of  $C^{-2}H$  bonds before and after a time period  $t_{\rm m}$ , the effect on the reorientation angle distribution  $W(\xi;t_{\rm m})$  produced by another isotropic reorientation process, but much slower than the rotational diffusion, is not detectable. Hence, its possible existence cannot be excluded. To detect other, slower, motional processes, the use of three- or higher-dimensional NMR techniques would be necessary.

# 5. 1D <sup>2</sup>H-NMR SPECTRA

To observe faster chain motion, on a time scale of  $10^{-4}$  to  $10^{-6}$  s, occurring at higher temperatures we obtained information from 1D  $^2$ H-NMR quadrupole echo spectra., namely at 1 bar (where  $T_{\rm g} = 250$  K) in the temperature range 280 to 315 K and at 4986 bar (where  $T_{\rm g} = 311$  K) for 348 and 353 K.



**Fig. 4.** Experimental quadrupole echo spectra at 284 and 290 at 1 bar and at 348 and 353 K at 4986 bar and the corresponding calculated spectra obtained from the discrete model (see text), as a function of the pulse distance.

Fig. 4 shows some quadrupole echo spectra as a function of the pulse distance  $\tau_1$ . Clearly, in this temperature range the line shape shows a strong dependence on the time scale of the molecular motion.

The effect of molecular motion on the line shape in quadrupole echo spectra can be described with a multisite exchange formalism<sup>17</sup>, that follows Abragam's treatment of motion narrowing<sup>18</sup>. With the help of a computer program, based on this formalism and kindly provided by the late Prof.Dr. Regitze Vold, we calculate the quadrupole echo spectrum resulting from the  $C^{-2}H$  bond jumping between N discrete orientations, using as a parameter the mean time  $\tau_j$  between jumps. In order to simulate the isotropic diffusive nature of the reorientation, we mimic it by an isotropic random walk on a sphere While the orientations of a  $C^{-2}H$  bond are limited to the orientations of the position vectors from the center to the carbon atoms in a  $C_{60}$  molecule. From every orientation only jumps to the three nearest neighbors are allowed. We again used the log-normal distribution (1) for  $\tau_j$ . Some of the results are shown in Fig. 4.

As can be seen, the discrete jump model results in satisfactory fits to spectra obtained at temperatures T=284 K and higher. The calculations do not yield acceptable fits to the spectra at lower temperatures. Presumably, this is due to the crudeness of the model; it breaks down when the correlation times become long and the reorientation angles become of the size corresponding to one jump in the discrete model. As in the calculations of the 2D spectra, the choice of asymmetric distributions does not result in substantially better fits.

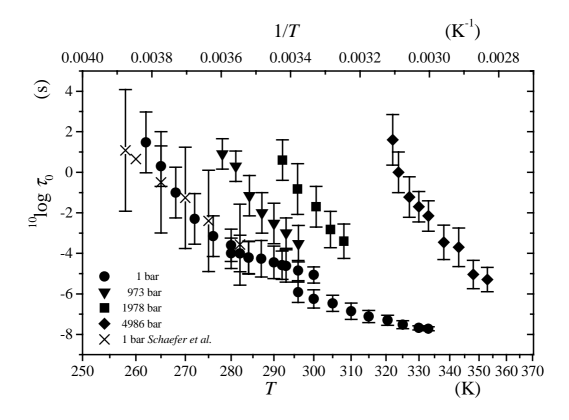
To relate  $\tau_j$  with  $\tau_d$ , and therefore  $\tau_{j0}$  with  $\tau_0$ , we have calculated the distribution  $W(\xi;t)$  of the relative reorientation angle  $\xi$  of a C-2H bond after a time t, for both the discrete model as described above and for isotropic diffusion, for which an analytical expression has been derived from the solution of the rotation diffusion equation (as presented in, e.g., reference<sup>19</sup>). From the comparison it appears that  $\tau_d \approx 4.5 \tau_j$ . The distributions of  $\tau_j$  are converted to distributions of  $\tau_d$  accordingly.

Information on still faster motion, on a time scale of  $10^{-6}$  to  $10^{-8}$  s, was derived from motional narrowed spectra<sup>18</sup>, obtained from free induction decays following a single  $\pi/2$  pulse. Above about  $T_{\rm g}+50$  K the motion of the C-<sup>2</sup>H bonds becomes sufficiently isotropic and rapid to average the EFG to zero, and the quadrupole echo pulse sequence fails to create a quadrupole echo. The experimental line shapes obtained at temperature values from  $T_{\rm g}+50$  to  $T_{\rm g}+83$  K (at ambient pressure only) are found to be non-Lorentzian. For

isotropic reorientation with a single value of the correlation  $\tau_d$  there is a simple relation <sup>18</sup> between the transverse relaxation rate  $!/T_2$  and  $\tau_d$ . Calculated line shapes are obtained by adding the Lorentzian line shapes, corresponding to values of  $\tau_d$ , and weighted again according to a log-normal distribution. In this way, good fits to the experimental spectra are obtained.

## 6. CHAIN MOTION AS A FUNCTION OF TEMPERATURE AND PRESSURE

All the parameters of the correlation time distributions, derived by fitting calculated spectra to the experimental spectra, are collected in Fig. 5. The indicated uncertainties in  $\tau_0$  and in  $\Delta_\sigma$  are estimates of the ranges of these parameters for which satisfactory fits are obtained. Along all four isobars the results show the characteristic slowing down of the dynamic process on approaching  $T_g$  from a higher temperature. At ambient pressure the time scale of the reorientation of the chain segments increases dramatically: from  $1.9 \cdot 10^{-8}$  s at 333 K to 30 s at 262 K, nine orders of magnitude in a temperature range of about



**Fig. 5.** The values of  $\tau_0$  (symbols) and  $\Delta_{\sigma}$  (vertical bars) of the correlation time distribution as a function of pressure as obtained by comparing experimental and calculated  ${}^2$ H-NMR spectra. Also included are the results by Schaefer et al  ${}^6$ .

70 K. The width of the correlation time distribution increases from 0.2 to 3.0 decades in the same temperature range. The results obtained at ambient pressure are in good agreement with the ones in the temperature range 300-370 K obtained from 2D exchange  $^2$ H NMR, reported in reference (also shown in Fig. 5). They also nicely connect with the ones obtained from  $^{13}$ C  $T_1$  data in the temperature range from 300 up to 370 K, where the correlation time is about  $2 \cdot 10^{-10}$  s. At higher pressure the behavior, although measured in smaller intervals of temperature, appears to be very similar.

As is well known, in many polymer materials the slowing down of the structural relaxation on approaching the temperature of the glass transition, may be described by the empirical WLF relation<sup>1,2</sup>. The form of this relation, in terms of an appropriate correlation time  $\tau(T)$ , is

$$\tau(T) = \tau(T_0) \exp[-c_1(T - T_0)/(T - T_0 + C_2)] \tag{4}$$

which is equivalent with the Vogel-Fulcher-Tammann-Hesse (VFTH) relation<sup>3,4,5</sup>

$$\tau(T) = \tau_1 \exp[C_0 / (T - T_V)] \tag{5}$$

with  $T_{\rm V}=T_0-C_2$ ,  $C_0=c_1C_2$  and  $\tau_1=\tau(T_0)\exp(-c_1)$ . In equation (4)  $T_0$  is a reference temperature. In fact, it appears that the temperature variation of the value of  $\tau_0$ , derived from our experiments can be described very well with the WLF-equation. We use it, for convenience, as a basis for the representation of the whole data set. By introducing  $C_1=2.303c_1$  one obtains the form which is usually found in the polymer literature:

$$^{10}\log(\tau_0(T)/\tau_0(T_g(p))) = -C_1(p)(T - T_g(p))/(C_2(p) + (T - T_g(p)))$$
 (6)

where  $T_0$  is set equal to the glass transition temperature  $T_g(p)$ .

We first used equation (6) to separately fit the data obtained at 1, 973, 1978 and 4986 bar. We reduce the number of free parameters by choosing for  $\tau_0(T_g(p))$  a constant value along the glass transition line, for which we take its average value  $10^{9.5}$  s as determined from separate fits to the data along isobars. The resulting WLF parameters  $C_1(p)$  and  $C_2(p)$  are collected in *Table 1*. They show no clear pressure dependence. The dependence of  $\tau_0$  on  $(T-T_g)$  does not change significantly at a pressure increase of up to 5 kbar, the

dependence of  $\tau_0$  on T and p can simply be expressed as  $\tau_0(T,p) = \tau_0(T-T_g(p))$ .

**Table I.** The parameters  $C_1(p)$  and  $C_2(p)$  resulting from fitting the temperature dependence of  $\tau_0$  at various pressure values with the WLF equation

p (bar)	$T_{\rm g}\left(p\right)  \left({ m K}\right)$	$C_1(p)$	$C_2(p)$ (K)
1.2±0.1	250±2	21.1±0.4	18.7±1.1
973±7	267±1	19.7±0.8	14.8±1.5
1978±14	279±2	20.3±0.9	16.7±1.7
4986±32	311±1	20.2±0.8	15.3±1.6

## 7. CHAIN MOTION AS A FUNCTION OF DENSITY

Since for pressure values up to 5 kbar the parameters  $C_1$  and  $C_2$  do not depend significantly on pressure, we can represent  $\tau_0(T,p)$  by an empirical WLF function, with  $< C_1>=20.7$  and  $< C_2>=16.4$ , the weighted averages of the values given in Table I:

$$^{10}\log(\tau_0(p,T)/10^{9.5}) = -20.7(T - T_g(p))/(16.4 + (T - T_g(p))). \tag{7}$$

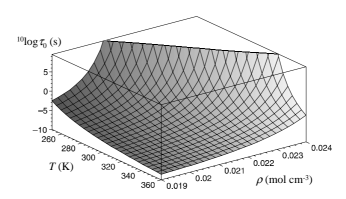
For aPP pVT-data are only available<sup>20,21,22</sup> for the melt. We have fitted the data of reference<sup>22</sup>, obtained in the range 295–582 K, 0–2000 bar for aPP with a weight average molar mass  $M_w$ =53400 g mol<sup>-1</sup> with the Tait equation<sup>23</sup>:

$$\rho(p,T) = \rho(0,T)[1 - D \ln[1 + p / E(T)]]^{-1}.$$
(8)

An accurate fit has been obtained by setting  $\rho(0,T)^{-1}=a_0+a_1T+a_2T^2$  with  $a_0=42.3858$  cm<sup>3</sup> mol<sup>-1</sup>,  $a_1=1.50521\cdot10^{-2}$  cm<sup>3</sup> mol<sup>-1</sup> K<sup>-1</sup> and  $a_2=2.83996\cdot10^{-5}$  cm<sup>3</sup> mol<sup>-1</sup> K<sup>-2</sup> and by taking  $E(T)=E_0\exp(-E_1T)$  with  $E_0=7256.3$  bar and  $E_1=5.0897\cdot10^{-3}$  K<sup>-1</sup>. The coefficient D is chosen to have the value 0.0894. We assume this equation-of-state to be also valid for our sample of aPP with molecular weight  $M_v\approx25000$  g mol<sup>-1</sup> and use it for an extrapolation of the pVT data to the glass-transition temperature and to 6000 bar.

By substituting equation (2), expressing the pressure along the glass-transition line as a function of  $T_g$ , in the EOS (9) we find that the density along the glass-transition line is represented by the quadratic function  $\rho(p_g) = b_0 + b_1 p_g + b_2 p_g^2$  with  $b_0 = 2.086 \cdot 10^{-2}$  molcm<sup>-3</sup>,  $b_1 = 6.400 \cdot 10^{-7}$  mol cm<sup>-3</sup> bar<sup>-1</sup> and  $b_2 = -3.789 \cdot 10^{-11}$  mol cm<sup>-3</sup> bar<sup>-2</sup>. A pressure increase of 5000 bar results in an increase of the density at the glass transition of about 11%. A very similar increase of the density along the glass-transition line has been observed in other

systems, for instance in poly(styrene)<sup>24</sup> (~2.1 % per kbar) and poly(carbonate)<sup>25</sup> (~2.0 % per kbar).



**Fig. 6.** WLF surface describing  $\tau_0$  as a function of temperature and density. The line at  $^{10}\log\tau_0=9.5$  indicates the glass transition.

By introducing the equation-of-state we obtain  $\tau_0$  as an empirical function of temperature and density. The resulting function is the surface shown in Fig. 6. Its intersection at  $^{10}\log\tau_0=9.5$  corresponds to the glass transition line. The figure reveals that along the paths at constant temperature and at constant density  $\tau_0$  shows the same steep increase resulting in the arrest of the chain motion at the glass transition.

The behavior of  $\tau_0$  along isothermals and along isochores is very similar. Over a large temperature and density range  $\tau_0$  shows a WLF-like dependence on  $T-T_g$  and on  $\rho(T_g)-\rho(T)$ , with parameters  $C_1$ ',  $C_2$ ' and  $C_1$ ",  $C_2$ " respectively. Again setting the value of  $\tau_0$  at the glass transition to  $10^{9.5}$ , we obtain an excellent fit of the isochores with the WLF equation, resulting in the values  $C_1$ '=20.5 and  $C_2$ '=19.8 K. These values reflect a weaker temperature dependence of  $\tau_0$  along isochores, in comparison with that of  $\tau_0$  along isobars, for which the  $C_1$  and  $C_2$  values are shown in Table I. The isothermals are equally

well fitted as a function of  $\rho(T_g)-\rho(T)$  by a WLF-like equation, with  $C_1$ "=20.1;  $C_2$ "=7.2·10<sup>-4</sup> mol cm<sup>-3</sup>. A fast WLF type increase in  $\tau_0$  occurs both on an isochoric and on an isothermal path to the glass transition. The temperature dependence of  $\tau_0$  along isobars is slightly stronger than along isochores. This effect is small due to the fact that the density variation along an isobar is only small, so that the temperature change rather than the volume change has the dominant effect on  $\tau_0$ .

As was noted above, the dependence of  $\tau_0$  on temperature and pressure can simply be expressed as  $\tau_0(T,p)=\tau_0(T-T_g(p))$ , the lines  $T-T_g(p)=$  constant are isochrones. Also at high pressure, the chain dynamics is completely determined by the same function of the temperature distance to the glass transition. This result is not self-evident, since the chain dynamics is strongly dependent on density. As can be derived from the equation-of state the density along lines of constant  $T-T_g(p)$  increases approximately in the same way as along the glass transition line, namely with about 11 % over 5 kbar. The decrease in the rate of motion by an increase in density is just compensated by the increase caused by raising the temperature to the value required to remain at constant  $T-T_g(p)$ .

# 8. THE RELATION OF THE WLF PARAMETERS WITH THERMODYNAMIC QUANTITIES

A number of theories of the glass transition offer a physical basis for this relation, in particular the statistical-mechanical theories of Adam and Gibbs<sup>26,27,28,29</sup> and of Parisi<sup>30</sup>, and the "free volume" theory<sup>2</sup>. We will compare our data with the results of the first theory in which the polymer melt is considered to be composed out of a number of subsystems, large enough to allow cooperative rearrangements of the chains into another configuration, without involving its environment. The subsystems are assumed to interact only weakly with the macroscopic system. It is shown, that the overwhelming majority of rearrangements occurs in the smallest regions, namely those which permit a rearrangement at all. The probability for a rearrangement in the system can be expressed in the ratio  $s_c/s_c$  of the configurational entropy of the smallest cooperatively rearranging region and that of the macroscopic system, and in the potential energy (per monomer segment)  $\Delta \mu$  hindering the cooperative rearrangement. This probability is proportional to  $\exp(-s_c\Delta \mu/s_c kT)$ . The configurational entropy of the smallest subsystem is  $s_c=k\ln(\Omega)$ , where  $\Omega$  is the associated minimum number of possible configurations. On lowering the temperature the smallest size of a cooperatively rearranging region will grow, until it

comprises the complete macroscopic system at some temperature  $T_2$ , and the polymer melt is effectively frozen. The configurational entropy of the macroscopic system vanishes at this temperature  $T_2$ , which is somewhat below the glass-transition temperature  $T_g$  determined in an experiment. With  $T_g$  chosen as the reference temperature, the WLF equation is derived with  $C_1$ =2.303 $\Delta\mu s_c/k\Delta C_pT_g\ln(T_g/T_2)$  and  $C_2$ = $T_g\ln(T_g/T_2)/[1+\ln(T_g/T_2)]$ .

We compare our values for  $C_1$  and  $C_2$  from Table 1 with the results of the Adam and Gibbs theory, and derive estimated values for  $T_2$  and  $\Delta\mu$ . The discontinuity in the specific heat is known to be  $\Delta C_p \approx 17.8 \text{ J mol}^{-1} \text{ K}^{-1}$  at ambient pressure<sup>31</sup>. In the range from ambient pressure to 5 kbar we have determined values for  $T_2$  from 231 to 295 K. On average  $T_g$ - $T_2$  $\approx$ 17 K, while from comparison with experiments Adam and Gibbs found  $T_g$ - $T_2$ =55±6 K for a number of glass-forming liquids at ambient pressure. With  $T_2$ =231 K we derive  $\Delta \mu s_c/k \approx 7.4$  kJ mol<sup>-1</sup> at ambient pressure. The rather high values for  $T_2$  are related to the relative low values found for  $C_2$  (when compared, for instance with the values found in reference  $^{26}$  for other substances). As the difference between  $T_{\rm g}$  and  $T_{\rm 2}$  is small, the result is quite sensitive to the factor  $ln(T_g/T_2)$ , and therefore also to the values used for  $T_2$ . To make a somewhat more quantitative comparison we choose  $\Omega=2$ , *i.e.* only two configurations are possible for the smallest cooperatively rearranging region, and find  $\Delta\mu \approx 5 \text{ kJ mol}^{-1}$  which is of the correct order of magnitude, namely that of the potential energy barrier between a trans and a gauche conformation of an alkane chain<sup>32</sup>. This is a reasonable result, further analysis would require the knowledge of  $\Delta C_p$  at high pressure. In principle, the Adam-Gibbs theory could be generalized to include the effect of density, but we did not attempt to do that.

Attempts to compare our data with the free volume theory<sup>2</sup> have lead to conflicting results. Therefore we concluded that a discussion of the effect of pressure on  $\tau_0$  in terms of the free volume theory is not very meaningful.

## 9. CONCLUSION

We studied the effect of high pressure on the motion of the main chain segments of aPP with different <sup>2</sup>H-NMR techniques in a large range of temperature above its glass transition temperature

At and above the glass transition small-angle motion of the chain segments, in which the chain C-<sup>2</sup>H bonds participate, is superimposed on the much slower quasi-isotropic rotational diffusive motion, which is the result of collective behavior. This process causes

the changes in the 1D and 2D  $^2$ H-NMR spectra. Clearly, this process may be identified with the  $\alpha$ -process observed in mechanical relaxation studies.

Our findings are consistent with the picture resulting from molecular dynamics simulations<sup>33</sup> of the local chain motion in amorphous polyethylene near the glass transition. The chain axis reorients much more slowly than a vector attached perpendicular to the chain axis. Fast small angle reorientation occurs within a restricted volume around the local main chain axis. In addition, chain segments take part in a slower quasi-isotropic rotational diffusion of the average local main chain axis, occurring as a result of the collective motion of its surroundings. On lowering the temperature the time scales of the two processes become unrelated. Eventually, the collective rearrangement of the surroundings of the chain segments becomes too slow to be detected on the time scale of the experiment. This arrest of the motion is the glass transition. The fast, small angle reorientation of the chain segments is not strongly affected by the glass transition.

From our experiments it follows that on approaching the glass transition the time scale of the collective motion of the main chain becomes longer very rapidly at decreasing temperature. The temperature dependence of the logarithmic average correlation time is shown to be very well described by a WLF- or VFTH-function. The collective motion of the main chain is strongly dependent on pressure (or rather on density). The main result of this investigation is that it shows that, on an isothermal approach to the glass transition, the chain dynamics as a function of density also follows a Vogel-Fulcher type law  $\tau(\rho) \propto \exp[C(\rho_{\rm VF}-\rho)^{-1}]$ , which, in our opinion, is an important experimental finding. Although this result is obtained by the use of a partially extrapolated equation-of-state, the errors introduced in the density can only be minor.

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